

Web Images Maps News Shopping Gmail more ▼



Sign in



compare core article using alignment algorithm


Search

Advanced Search  
Preferences

Web: Result details: SML

Results 1 - 10 of about 327,000 for **compare core article using alignment algorithm**. (0.22 seconds)

## Scholarly articles for **compare core article using alignment algorithm**

-  [Using 3D Hidden Markov Models that explicitly represent ...](#) - Alexandrov - Cited by 19
- [A comprehensive comparison of multiple sequence ...](#) - Thompson - Cited by 318
- [Analysis of covariation in an SH3 domain sequence ...](#) - Larson - Cited by 62

### 1. A new progressive-iterative **algorithm** for multiple structure ...

As a result, a variety of multiple structure **alignment algorithms** have .... (3.4 )  
Minimize the RMS fluctuation of the **core** by **using** a SIMPLEX optimization. ...  
[bioinformatics.oxfordjournals.org/cgi/content/full/21/15/3255](#) - [Similar pages](#)  
by D Lupyán - 2005 - [Cited by 35](#) - [Related articles](#)

### 2. Statistical evaluation and comparison of a pairwise **alignment** ...

In this comparison, SANK\_AL performed sequence **alignment** allowing from 1 to 25  
gaps. Figures 2 and 3 show the results of each **algorithm using** BLOSUM62, ...  
[bioinformatics.oxfordjournals.org/cgi/content/full/21/8/1421](#) - [Similar pages](#)  
by Y Nozaki - 2005 - [Cited by 3](#) - [Related articles](#) - [All 6 versions](#)  
[More results from bioinformatics.oxfordjournals.org »](#)

### 3. PLoS Computational Biology: Recent Evolutions of Multiple Sequence ...

The scoring schemes used by the pairwise **alignment algorithm** are arguably .... Find  
this **article** online; Taylor WR (1999) Protein structure comparison **using** ...  
[www.ploscompbiol.org/article/info:doi/10.1371/journal.pcbi.0030123](#) - 74k  
- [Cached](#) - [Similar pages](#)  
by C Notredame - 2007 - [Cited by 9](#) - [Related articles](#) - [All 8 versions](#)

### 4. Fast **algorithms** for large-scale genome **alignment** and comparison

This **article** has been cited by other **articles** in PMC. ... This **core algorithm** is  
implemented **using** a suffix-tree data structure, which permits very fast and ...  
[www.pubmedcentral.nih.gov/articlerender.fcgi?artid=117189](#) - [Similar pages](#)  
by AL Delcher - 2002 - [Cited by 245](#) - [Related articles](#) - [All 38 versions](#)

### 5. A novel approach to structural **alignment** using realistic ...

Here we present a novel approach to structural **alignment using** a clique finding **algorithm** and environmental information (SAUCE). In this approach, we build ...  
[www.proteinscience.org/cgi/content/abstract/14/12/2935](http://www.proteinscience.org/cgi/content/abstract/14/12/2935) - [Similar pages](#)  
by Y Chen - 2005 - [Cited by 6](#) - [Related articles](#) - [All 7 versions](#)

6. **Smith-Waterman algorithm** - Wikipedia, the free encyclopedia

Local **alignment** avoids such regions altogether and focuses on those with a ... In 2000, a fast implementation of the Smith-Waterman **algorithm using** the SIMD ...  
[en.wikipedia.org/wiki/Smith-Waterman](http://en.wikipedia.org/wiki/Smith-Waterman) - 42k - [Cached](#) - [Similar pages](#)

7. **Algorithms** for Molecular Biology | Full text | Multiple sequence ...

Most multi-**alignment** programs are **using** heuristic optimisation **algorithms**, i.e. they are, in general, not able to find the mathematically optimal **alignment** ...  
[www.almob.org/content/1/1/6](http://www.almob.org/content/1/1/6) - 111k - [Cached](#) - [Similar pages](#)  
by B Morgenstern - 2006 - [Cited by 14](#) - [Related articles](#)

8. **MUSTANG: A multiple structural alignment algorithm**

In this **article**, we present an **algorithm** for multiple ..... sequence **alignment using** ClustalW, and structural **alignment using** MUSTANG, of the two odorant- ...  
[doi.wiley.com/10.1002/prot.20921](http://doi.wiley.com/10.1002/prot.20921) - [Similar pages](#)  
by AS Konagurthu - 2006 - [Cited by 51](#) - [Related articles](#) - [All 5 versions](#)

9. **Fast algorithms** for large scale genome **alignment** and comparison

Fast **algorithms** for large scale genome **alignment** and comparison. From mala, 8 months ago. A presentation of the three **articles** about MUMmer by A.L. Delcher ...  
[www.slideshare.net/mala/fast-algorithms-for-large-scale-genome-alignment-and-comparison/](http://www.slideshare.net/mala/fast-algorithms-for-large-scale-genome-alignment-and-comparison/) - 88k - [Cached](#) - [Similar pages](#)

10. **BioMed Central** | Full text | TOPS++FATCAT: fast flexible ...

TOPS++FATCAT: Fast flexible structural **alignment using** constraints derived ... In the second approach, specialized comparison **algorithms**, such as DALI [3], ...  
[www.biomedcentral.com/1471-2105/9/358](http://www.biomedcentral.com/1471-2105/9/358) - 83k - [Cached](#) - [Similar pages](#)  
by M Veeramalai - 2008

Google    
1 2 3 4 5 6 7 8 9 10 [Next](#)

[Search within results](#) | [Language Tools](#) | [Search Tips](#) | [Dissatisfied? Help us improve](#) | [Try Google Experimental](#)

---

[Google Home](#) - [Advertising Programs](#) - [Business Solutions](#) - [Privacy](#) - [About Google](#)